

## **Training-specific functional, neural, and hypertrophic adaptations to explosive- vs. sustained-contraction strength training**

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**TITLE PAGE**

**Title:**

TRAINING SPECIFIC FUNCTIONAL, NEURAL AND HYPERTROPHIC  
ADAPTATIONS TO EXPLOSIVE- VS. SUSTAINED-CONTRACTION STRENGTH  
TRAINING

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**Abbreviated title for running head:**

EXPLOSIVE- VS. SUSTAINED-CONTRACTION STRENGTH TRAINING

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Resistance exercise; Neural drive; Rate of torque development; Maximum strength;  
Contractile properties

## Abstract

Training specificity is considered important for strength training, although the functional and underpinning physiological adaptations to different types of training, including brief explosive contractions, are poorly understood. This study compared the effects of 12-wks of explosive-contraction (ECT,  $n=13$ ) vs. sustained-contraction (SCT,  $n=16$ ) strength training vs. control (CON,  $n=14$ ) on the functional, neural, hypertrophic, and intrinsic contractile characteristics of healthy young men. Training involved 40 isometric knee extension repetitions (x3/wk): contracting as fast and hard as possible for ~1 s (ECT); or gradually increasing to 75% of maximum voluntary torque (MVT) before holding for 3 s (SCT). Torque and EMG during maximum and explosive contractions, torque during evoked octet contractions, and total quadriceps muscle volume (QUADS<sub>VOL</sub>) were quantified pre and post training. MVT increased more after SCT than ECT (23 vs. 17%; effect size [ES]=0.69), with similar increases in neural drive, but greater QUADS<sub>VOL</sub> changes after SCT (8.1 vs. 2.6%; ES=0.74). ECT improved explosive torque at all time points (17-34%;  $0.54 \leq ES \leq 0.76$ ) due to increased neural drive (17-28%), whereas only late-phase explosive torque (150 ms, 12%; ES=1.48) and corresponding neural drive (18%) increased after SCT. Changes in evoked torque indicated slowing of the contractile properties of the muscle-tendon unit after both training interventions. These results showed training-specific functional changes that appeared to be due to distinct neural and hypertrophic adaptations. ECT produced a wider range of functional adaptations than SCT, and given the lesser demands of ECT this type of training provides a highly efficient means of increasing function.

## **New & Noteworthy**

Explosive-contraction strength training (ECT) denoted by brief contractions with high rate of torque development produced a wider range of functional adaptations than sustained-contraction strength training (SCT), with improvements in early- and late-phase explosive strength, as well as maximum strength. In contrast, SCT only improved maximum and late-phase explosive strength. The substantially lower loading duration of ECT (7% of SCT) makes this a less-demanding training modality compared to SCT, which may be preferentially tolerated by musculoskeletal patients.

## **Introduction**

Maximum and explosive strength are two components of skeletal muscle function that can be critical to the performance of human movement. Maximum strength is the greatest amount of force that can be generated, whereas explosive strength reflects the ability to increase force rapidly from a low or resting level (1, 20, 48). Muscle weakness, including low maximum and explosive strength, contributes to the functional limitations experienced by numerous patient groups (35, 42, 43), including osteoarthritis patients (26). Strength training is widely recommended for improving function of all adults (32, 39) and the increases in explosive and/or maximum strength that occur following training may have profound benefits to mobility, locomotion, and quality of life of older individuals and patients (13, 25, 26, 34, 37, 44). Whilst training specificity is widely considered important within the context of strength training (8, 12, 18, 21), the functional adaptations to different types of strength training are not well understood, reducing the efficacy of training guidance and prescription. Furthermore, the similarity or specificity of the underpinning neural and contractile adaptations to different training regimes has received relatively little attention.

80

81 Explosive-contraction strength training (ECT), emphasizing rapid torque development during  
82 short contractions, is a relatively non-fatiguing training modality that may be well tolerated  
83 by patient groups (i.e. osteoarthritis) who commonly report substantial fatigue (36, 38) and  
84 therefore may offer improved adherence within these populations. ECT has been found to  
85 produce significant increases in both maximum and explosive strength (48). In contrast,  
86 conventional strength training typically has a primary emphasis of training with sustained-  
87 contractions (SCT) at high loads leading to pronounced fatigue (31) and may neglect rapid  
88 torque development. Our recent 4-week intervention study contrasted ECT and SCT finding  
89 distinct training-specific adaptations in functional capabilities and neural drive: maximum  
90 strength and corresponding electromyography (EMG) increased more after SCT; and early-  
91 phase explosive strength and EMG ( $\leq 100$  ms) during the rising/explosive phase of  
92 contraction increased more following ECT (45). This demonstrated that at least in the initial  
93 stages of a training programme ECT and SCT produce distinct functional and neural  
94 adaptations. However the efficacy of longer-term ECT for functional and neural adaptations  
95 remains unknown, and the contrasting influence of these training interventions on the  
96 intrinsic contractile properties and hypertrophy (volume) of skeletal muscle has yet to be  
97 investigated. A more comprehensive comparison of ECT and SCT may facilitate a greater  
98 understanding of the influence of training variables, particularly loading duration (high SCT  
99 vs. low ECT) and rate of torque development (RTD, high ECT vs. low SCT), on functional  
100 and physiological adaptations.

101

102 The purpose of this study was to investigate the efficacy of 12 weeks of ECT, and compare it  
103 to SCT and a control group (CON) by assessing the specificity of the functional changes  
104 (maximum and explosive strength), as well as the underpinning adaptations in neural drive,

intrinsic contractile properties, and muscle volume after these interventions. We hypothesised that ECT and SCT would elicit distinct and specific functional changes (ECT>SCT for early-phase [ $\leq 100$  ms] explosive strength; SCT>ECT for maximum strength), as a result of distinct neural and contractile adaptations.

## **Materials and Methods**

### *Participants*

Forty-eight young, healthy, asymptomatic, males who had not completed lower-body strength training for >18 months and were not involved in systematic physical training were recruited and provided written informed consent prior to participation in this study that was approved by the Loughborough University Ethical Advisory Committee. Following familiarization participants were randomly assigned to ECT, SCT, or CON groups that were matched for maximum voluntary torque (MVT) and body mass. A total of five participants withdrew from the study (four due to personal reasons and one was excluded due to non-compliance). Forty-three participants (ECT [ $n=13$ ]; SCT [ $n=16$ ]; CON [ $n=14$ ]) completed the study. Baseline recreational physical activity was assessed with the International Physical Activity Questionnaire (IPAQ, short format (14)).

### *Overview*

Participants visited the laboratory for a familiarisation session involving voluntary maximum and explosive, as well as evoked twitch contractions to facilitate group allocation. Thereafter, two duplicate laboratory measurement sessions were conducted both pre (sessions 7-10 days apart prior to the first training session) and post (2-3 days after the last training session and 2-

3 days later) 12-weeks of unilateral knee extensor strength training. Axial T1-weighted MRI scans of the thigh were also conducted pre (5 days prior to the first training session) and post (2-3 days after the final training session). Training and testing were completed with the same isometric apparatus. Training for the ECT and SCT groups involved unilateral isometric contractions of both legs three times a week for 12-weeks (36 sessions in total), whereas CON participants attended only the measurement sessions and maintained their habitual activity. All participants were instructed to maintain their habitual physical activity and diet throughout the study. Laboratory testing sessions involved recordings of the dominant leg isometric knee-extension torque and surface EMG of the superficial quadriceps muscles during voluntary maximum and explosive contractions, as well as evoked maximum twitch and octet contractions (via electrical stimulation of the femoral nerve). Measurement sessions were at a consistent time of day and started between 12:00-19:00.

### *Training*

After a brief warm-up of sub-maximum contractions of both legs, participants completed four sets of ten unilateral isometric knee-extensor contractions of each leg; with sets alternating between dominant and non-dominant legs until 4 sets per leg had been completed. Each set took 60 s with 2 min between successive sets on the same leg. ECT involved short, explosive contractions with participants instructed to perform each contraction “as fast and hard as possible” up to  $\geq 80\%$  MVT for  $\sim 1$  s, and then relax for 5 s between repetitions (Fig. 1 A). A computer monitor displayed RTD (10-ms time epoch) to provide biofeedback of explosive performance, with a cursor indicating the highest peak RTD achieved throughout the session, participants were encouraged to achieve a higher peak RTD with each subsequent contraction. The torque-time curve was also shown: firstly, with a horizontal cursor at 80% MVT (target

force) to ensure sufficiently forceful contractions, and secondly, on a sensitive scale highlighting baseline torque in order to observe and correct any pre-tension or countermovement.

SCT involved sustained contractions at 75%MVT, with 2 s rest between contractions. In order to control the RTD these participants were presented with a target torque trace 2 s before every contraction and instructed to match this target, which increased torque linearly from rest to 75%MVT over 1 s before holding a plateau at 75%MVT for a further 3 s. All training participants (ECT and SCT) performed three maximum voluntary isometric contractions (MVCs, see below) at the start of each training week in order to re-establish MVT and prescribe training torques. Torque data from the first training session of weeks 1, 6 and 12 were analysed for all training participants (i.e. ECT and SCT) in order to quantify peak loading magnitude (peak torque, mean of all repetitions), loading rate (peak RTD, 50-ms epoch, mean of all repetitions), and loading duration (defined as time >65%MVT per session).

#### *Force and EMG recording*

Measurement and training sessions were completed in a rigid custom-made isometric dynamometer with knee and hip angles of 115° and 126° (180° = full extension), respectively. Adjustable straps were tightly fastened across the pelvis and shoulders to prevent extraneous movement. An ankle strap (35 mm width reinforced canvas webbing) was placed ~15% of tibial length (distance from lateral malleolus to knee joint space), above the medial malleolus, and positioned perpendicular to the tibia and in series with a calibrated S-beam strain gauge (Force Logic, Swallowfield, UK). The analogue force signal from the strain gauge was amplified (x370) and sampled at 2,000 Hz using an external A/D converter (Micro 1401;



CED Ltd., Cambridge, UK) and recorded with Spike 2 computer software (CED Ltd., Cambridge, UK). In offline analysis, force data were low-pass filtered at 500 Hz using a fourth-order zero-lag Butterworth filter (33), gravity corrected by subtracting baseline force, and multiplied by lever length, the distance from the knee joint space to the centre of the ankle strap, to calculate torque values.

Surface EMG was recorded from the superficial quadriceps muscles (rectus femoris [RF], vastus lateralis [VL], vastus medialis [VM]) using a wireless EMG system (Trigno; Delsys Inc., Boston, MA). Following skin preparation (shaving, abrading, and cleansing with 70% ethanol), Single differential Trigno Standard EMG sensors (Delsys Inc., Boston, MA) each with a fixed 1 cm inter-electrode distance were attached at six separate sites over the superficial quadriceps muscles at set percentages of thigh length above the superior border of the patella (RF 65 and 55%; VL 60 and 55%; VM 35 and 30%) and parallel to the presumed orientation of the underlying fibres. EMG signals were amplified at source (x300; 20- to 450-Hz bandwidth) before further amplification (overall effective gain, x909), and sampled at 2,000 Hz via the same A/D converter and computer software as the force signal, to enable data synchronization. In offline analysis, EMG signals were corrected for the 48-ms delay inherent to the Trigno EMG system and band-pass filtered (6-500 Hz) using a fourth-order zero-lag Butterworth filter.

#### *Pre and post measurement sessions*

Following a brief warm-up of the dominant leg (3 s contractions at 50% [x3], 75% [x3], and 90% [x1] of perceived maximum) measurements were completed in the following order.

200

201 *Maximum voluntary contractions*

202 Participants performed 3-4 MVCs and were instructed to “push as hard as possible” for 3-5 s  
203 and rest for  $\geq 30$  s between efforts. A torque-time curve with a horizontal cursor indicating the  
204 greatest torque obtained within that session was displayed for biofeedback and verbal  
205 encouragement was provided during all MVCs. Knee extensor MVT was the greatest  
206 instantaneous torque achieved during any MVC or explosive contraction during that  
207 measurement session. Root mean square (RMS) EMG for a 500 ms epoch at MVT (250 ms  
208 either side) was calculated for each electrode site before averaging across the six sites to  
209 provide a whole quadriceps measurement ( $QEMG_{MVT}$ ). In addition, RMS EMG at MVT was  
210 normalized to  $M_{MAX}$  area (see below) from the corresponding EMG electrode site and then  
211 averaged across all quadriceps EMG sites.

212

213 *Explosive voluntary contractions*

214 Participants completed ten explosive voluntary contractions. They were instructed to perform  
215 each contraction “as fast and hard as possible” for  $\sim 1$  s, in order to exceed 80%MVT, and  
216 then relax for  $\geq 15$  s between contractions. Contractions with a change in baseline torque (pre-  
217 tension or countermovement) of  $>0.34$  Nm in the 300 ms prior to contraction onset were  
218 discarded. The three best contractions (highest torque at 100 ms) were analysed in detail for  
219 torque and EMG. Voluntary explosive torque was measured at 50, 100, and 150 ms from  
220 contraction onset ( $T_{50}$ ,  $T_{100}$ , and  $T_{150}$ ), before averaging across the three contractions.  
221 Explosive torque was also expressed relative to MVT to assess if explosive and maximum  
222 strength changed proportionally.

223

224 RMS EMG of each of the quadriceps sensor sites was measured over three time periods: 0-50,  
225 0-100 and 0-150 ms from EMG onset of the first agonist muscle to be activated (see below),  
226 before averaging to produce overall quadriceps measurements ( $QEMG_{0-50}$ ,  $QEMG_{0-100}$ ,  
227  $QEMG_{0-150}$ ) for the three best contractions. RMS EMG values from each sensor were also  
228 normalized to both  $EMG_{MVT}$  and  $M_{MAX}$  area for that site before averaging. To decide whether  
229 to report absolute RMS EMG or RMS EMG normalized to  $M_{MAX}$  the intra-participant  
230 reproducibility of  $EMG_{MVT}$  for both EMG measures was assessed over the 12-week  
231 intervention for CON (see below), and the most reproducible measure used. The ratio of  
232 Voluntary  $T_{50}$ /Octet  $T_{50}$  (see below) was used as an additional measure of volitional neural  
233 efficacy during the voluntary explosive contractions.

234

235 During offline analysis, all torque and EMG onsets were identified manually by visual  
236 identification by one trained investigator using a systematic approach (46, 49) considered to  
237 be more valid than automated methods (49). Briefly, torque and EMG signals were initially  
238 viewed on an  $x$  axis scale of 300 ms prior to the contraction and  $y$  axis scales of 0.68 Nm  
239 (torque) or 0.05 mV (EMG) (46, 49) before zooming in to determine the instant of the last  
240 peak or trough before the signal deflected away from the envelope of the baseline noise.

241

#### 242 *Evoked twitch and octet contractions*

243 A constant current variable voltage stimulator (DS7AH; Digitimer Ltd., Welwyn Garden City,  
244 UK), cathode probe (1-cm diameter, Electro-Medical Supplies Ltd., Wantage, UK), and  
245 anode electrode (7 x 10 cm carbon rubber electrode; Electro-Medical Supplies Ltd., Wantage,  
246 UK) were used to electrically stimulate the femoral nerve. The cathode and anode were  
247 coated with electrode gel and securely taped to the skin over the femoral nerve in the femoral

triangle and over the greater trochanter, respectively. Cathode location was determined by delivering single electrical impulses (square wave-pulses of 0.2 ms duration,  $\geq 12$  s apart) in order to identify the position that elicited the greatest sub-maximum twitch response. The current intensity was progressively increased until plateaus in peak twitch force and peak-to-peak M-wave amplitude were reached. Then three supra-maximal twitch and  $M_{MAX}$  responses were evoked (15 s apart) at a higher current ( $\geq 50\%$ ) to ensure supra-maximal stimulation. The following variables were averaged across the three supra-maximal twitch contractions: peak twitch torque (Twitch Peak T); absolute torque (Twitch  $T_{50}$ ) and torque expressed relative to Twitch Peak T (Relative Twitch  $T_{50}$ ) at 50 ms after contraction onset; time from contraction onset to peak twitch torque (Twitch TPT); and the cumulative  $M_{MAX}$  area from EMG onset to the point where the signal returned to baseline for each of the six EMG sites.

During the second pre and first post measurement sessions only, octet contractions (eight impulses at 300 Hz) were evoked at progressive currents ( $\geq 15$  s apart) until a plateau in the amplitudes of peak torque and peak RTD were achieved. Then, three discrete pulse trains ( $\geq 15$  s apart) were delivered with a higher current ( $\geq 20\%$  to ensure supra-maximal stimulation) to evoke maximum octet contractions. Peak torque (Octet Peak T), absolute torque (Octet  $T_{50}$ ) and torque expressed relative to Octet Peak T (Relative Octet  $T_{50}$ ) at 50 ms after contraction onset, and time from contraction onset to Octet Peak T (Octet TPT) were averaged across the three maximum octet contractions. Due to the discomfort caused by the octet contractions a total of seven participants across the three groups were unable to tolerate this measurement.

272

273 *Muscle volume*

274 A 1.5T MRI scan of the dominant leg was made in the supine position at a knee joint angle of  
275  $\sim 163^\circ$  using a receiver 8-channel whole body coil (Signa HDxt, GE). T1-weighted axial  
276 slices (5 mm thick, 0 mm gap) were acquired from the anterior superior iliac spine to the knee  
277 joint space in two overlapping blocks. Oil filled capsules placed on the lateral side of the  
278 participants' thigh allowed alignment of the blocks during analysis. MR images were  
279 analyzed by two investigators using Osirix software (version 6.0, Pixmeo, Geneva,  
280 Switzerland). Pre and post scans of each participant were analyzed by the same investigator.  
281 The quadriceps (RF, VL, VM, and vastus intermedius; VI) muscles were manually outlined  
282 in every third image (i.e. every 15 mm) starting from the most proximal image in which the  
283 muscle appeared. The volume of each muscle was calculated using cubic spline interpolation  
284 (GraphPad Prism 6, GraphPad Software, Inc.). Total quadriceps volume ( $QUADS_{VOL}$ ) was  
285 the sum of the individual muscle volumes. Inter- and intra-rater reliability for  $QUADS_{VOL}$   
286 calculated from the repeated analysis of five MRI scans was 1.2% and 0.4%, respectively.  
287 Data from one participant was excluded due to excessive movement artifacts.

288

289 *Data analysis and statistics*

290 All data was anonymized prior to analysis. Reproducibility of the measurements over the 12-  
291 week intervention period was calculated for CON (pre vs. post) as within-participant  
292 coefficient of variation ( $CV_W$ ;  $(SD/mean) \times 100$ ). MVT and  $QEMG_{MVT}$  measurements from  
293 the duplicate test sessions were averaged to produce criterion pre and post values for  
294 statistical analysis; unless the  $CV_W$  for the MVT was  $\geq 10\%$  (calculated from duplicate test  
295 sessions), in which case the lowest MVT value and corresponding  $QEMG_{MVT}$  were discarded.  
296 Mean  $T_{50}$ ,  $T_{100}$ , and  $T_{150}$  and corresponding QEMG ( $QEMG_{0-50}$ ,  $QEMG_{0-100}$ ,  $QEMG_{0-150}$ )

from the duplicate test sessions were used as criterion pre and post values for statistical analysis; unless the  $CV_W$ , (calculated from duplicate test sessions at the given time point) for  $T_{50}$  was  $\geq 20\%$ , in which case a weighted mean for all three explosive torque time points and corresponding QEMG measures were used.

All statistical analyses were performed using SPSS Version 22.0 (IBM Corp., Armonk, NY). Data are reported as means  $\pm$  SD; apart from within figures where data are mean  $\pm$  standard error of the mean (SE) for presentation purposes. One-way ANOVAs were conducted on all pre-test variables to assess whether baseline differences existed between groups. Unpaired  $t$ -tests were used to assess differences in training variables (loading rate, duration, and magnitude) between ECT and SCT. Within-group changes were evaluated with paired  $t$ -tests. Comparison of between-group adaptations to the intervention were assessed with repeated measures analysis of co-variance (ANCOVA; group [ECT vs. SCT vs. CON] x time [pre vs. post]), with corresponding pre training values used as covariates. When group x time interaction effects displayed  $P < 0.05$  then post-hoc tests were conducted and included the calculation of effect size (ES) and least significant differences (LSD) of absolute changes (pre to post) between groups (i.e., ECT vs. SCT, ECT vs. CON, and SCT vs. CON). ES for absolute change data was calculated as previously detailed for between-subject study designs [30] and classified as:  $< 0.20 =$  “trivial”;  $0.20-0.50 =$  “small”;  $0.50-0.80 =$  “moderate”; or  $> 0.80 =$  “large”. Least significant difference (LSD) post-hoc tests were produced from one-way ANCOVAs and were corrected for multiple comparisons (5). We considered there to be good evidence of between-group differences if both  $ES > 0.50$  and LSD post-hoc  $P < 0.10$ .

## Results

### *Group characteristics at baseline*

At baseline no differences (ANOVA,  $P \geq 0.767$ ) were observed between groups for habitual physical activity (IPAQ: ECT  $2047 \pm 1081$ ; SCT  $2135 \pm 1230$ ; CON  $2321 \pm 1614$  metabolic equivalent min/wk), age (ECT  $25 \pm 2$ ; SCT  $25 \pm 2$ ; CON  $25 \pm 3$  yr), body mass (ECT  $70 \pm 10$ ; SCT  $71 \pm 9$ ; CON  $72 \pm 7$  kg), or height (ECT  $1.74 \pm 0.07$ ; SCT  $1.75 \pm 0.08$ , CON  $1.76 \pm 0.06$  m). Similarly, no baseline differences were detected for functional, neural, intrinsic contractile properties, or muscle volume.

### *Reproducibility of Torque and EMG measurements*

The reproducibility of pre and post measures for the CON group over the 12-week period was excellent for MVT,  $T_{100}$ , and  $T_{150}$  ( $CV_W$  2.9, 4.4 and 4.9% respectively), but poor for  $T_{50}$  ( $CV_W$  15.7%). Absolute  $EMG_{MVT}$  (9.8%) had better  $CV_W$  than  $EMG_{MVT}$  normalized to  $M_{MAX}$  area (14.7%), and therefore, absolute EMG data are presented. Twitch (Twitch  $T_{50}$ , Twitch Peak T, Relative Twitch  $T_{50}$ , Twitch TPT) and octet ( $n=11$ , Octet  $T_{50}$ , Octet Peak T, Relative Octet  $T_{50}$ , Octet TPT) variables displayed excellent to good  $CV_W$  values (1.8-6.1%).

### *Training quantification for ECT vs. SCT*

Loading duration, quantified as time  $>65\%$  MVT per session, was greater for SCT than ECT (unpaired  $t$ -test  $P < 0.001$ ; Fig. 1 B). Conversely, ECT involved  $\sim 6$ -fold greater RTD per repetition than SCT (unpaired  $t$ -test  $P < 0.001$ ; Fig. 1 C). Peak loading magnitude was also slightly greater for ECT than SCT ( $81 \pm 4$  vs.  $75 \pm 2\%$  MVT; unpaired  $t$ -test  $P < 0.001$ ).

### *Voluntary torque*

MVT increased after ECT and SCT (both paired *t*-test  $P < 0.001$ ), but not following CON ( $P = 0.739$ ; Table 1 & 4). The absolute increase in MVT was greater than CON for both ECT and SCT (both  $ES \geq 2.06$  “large”, LSD  $P < 0.001$ ), and 38% larger after SCT than ECT ( $ES = 0.69$  “moderate”,  $P = 0.052$ ; Fig. 2).

Explosive torque increased at  $T_{50}$ ,  $T_{100}$ , and  $T_{150}$  after ECT (paired *t*-test  $P = 0.047$ ,  $P = 0.008$ , and  $P < 0.001$ , respectively; Table 1 & 4). Whereas, there were no changes in explosive torque after CON (paired *t*-test  $0.420 \leq P \leq 0.847$ ) and only  $T_{150}$  increased following SCT ( $P < 0.001$ ) with no change in  $T_{50}$  or  $T_{100}$  ( $0.140 \leq P \leq 0.939$ ). Group comparisons revealed that ECT produced greater increases in explosive torque than SCT after 100 ms, but not after 150 ms ( $T_{100}$ :  $ES = 0.72$  “moderate”, LSD  $P = 0.092$ ;  $T_{150}$ :  $ES = 0.54$  “moderate”,  $P = 0.145$ ), and larger increases than CON from 100 ms onwards ( $T_{100}$ :  $ES = 0.98$  “large”,  $P = 0.042$ ;  $T_{150}$ :  $ES = 1.59$  “large”,  $P < 0.001$ ). SCT resulted in greater increases than CON only at  $T_{150}$  ( $ES = 1.48$  “large”, LSD  $P = 0.008$ ).

Relative explosive torque (%MVT), at all time points, decreased following SCT (paired *t*-test  $0.004 \leq P \leq 0.032$ ; Table 1), but remained unchanged after ECT and CON ( $0.344 \leq P \leq 0.984$ ). The decrease in relative explosive torque after SCT was greater than ECT ( $T_{100}$ :  $ES = 0.88$  “large”, LSD  $P = 0.015$ ; and  $T_{150}$ :  $ES = 0.91$  “large”,  $P = 0.006$ ) and CON ( $T_{100}$ :  $ES = 1.15$  “large”,  $P = 0.016$ ; and  $T_{150}$ :  $ES = 0.99$  “large”,  $P = 0.022$ ; Fig. 3). Changes in relative explosive torque did not differ between ECT and CON ( $T_{100}$ :  $ES = 0.11$  “trivial”, LSD  $P = 0.844$ ; and  $T_{150}$ :  $ES = 0.12$  “trivial”,  $P = 0.547$ ).



*Neural drive*

QEMG<sub>MVT</sub> increased, or had a tendency to increase, after SCT (paired *t*-test  $P < 0.001$ ) and ECT ( $P = 0.099$ ), but not CON ( $P = 0.130$ ; Table 2 and 4). The increase in QEMG<sub>MVT</sub> was greater than CON for both ECT (ES=0.87 “large”, LSD  $P = 0.018$ ) and SCT (ES=2.30 “large”,  $P < 0.001$ ), but was not different between ECT and SCT (ES=0.36 “small”,  $P = 0.370$ ; Fig. 2). QEMG<sub>0-50</sub>, QEMG<sub>0-100</sub>, and QEMG<sub>0-150</sub> increased or had a tendency to increase after ECT (paired *t*-test  $P = 0.089$ ,  $P = 0.048$  and  $P = 0.003$ , respectively; Table 2). There were no changes in explosive QEMG measurements after CON, and only QEMG<sub>0-150</sub> increased after SCT (paired *t*-test  $P = 0.009$ ; Table 2). Group comparisons showed ECT to increase explosive neural drive by more than CON at all time points (QEMG<sub>0-50</sub>: ES=0.85 “large”, LSD  $P = 0.036$ ; QEMG<sub>0-100</sub>: ES=1.07 “large”,  $P = 0.018$ ; QEMG<sub>0-150</sub>: ES=1.57 “large”,  $P < 0.001$ ; Fig. 3), and by more than SCT for QEMG<sub>0-150</sub> (ES=0.58 “moderate”,  $P = 0.061$ ) but not earlier periods (QEMG<sub>0-50</sub>: ES=0.58 “moderate”,  $P = 0.101$ ; or QEMG<sub>0-100</sub>: ES=0.46 “small”,  $P = 0.254$ ; Fig. 3). SCT increased QEMG<sub>0-150</sub> more than CON (ES=1.20 “large”, LSD  $P = 0.021$ ), but this was not the case for earlier periods ( $0.30 \leq ES \leq 0.61$  “small” to “moderate”,  $0.154 \leq P \leq 0.463$ ).

Relative explosive neural drive (as % QEMG<sub>MVT</sub>) for all time periods decreased after SCT (paired *t*-test,  $0.001 \leq P \leq 0.004$ ), but not following ECT or CON ( $P \geq 0.395$ ; Table 2). After SCT the decreases in relative QEMG<sub>0-100</sub> were greater than ECT (ES=0.59 “moderate”, LSD  $P = 0.086$ ) and CON (ES=0.99 “large”,  $P = 0.045$ ), as was QEMG<sub>0-150</sub> vs. ECT (ES=0.62 “moderate”,  $P = 0.066$ ). Changes in relative explosive QEMG<sub>0-100</sub> and QEMG<sub>0-150</sub> did not differ between ECT and CON ( $0.02 \leq ES \leq 0.29$ , LSD  $0.623 \leq P \leq 0.697$ )

Voluntary T<sub>50</sub>/Octet T<sub>50</sub> ratio appeared to increase after ECT ( $n=12$ ; pre  $42 \pm 20\%$  vs. post  $53 \pm 19\%$ ) but did not reach statistical significance for the within group change (paired  $t$ -test  $P=0.122$ ) or group  $\times$  time interaction effect (ANCOVA,  $P=0.107$ ). No changes in the Voluntary T<sub>50</sub>/Octet T<sub>50</sub> ratio occurred after SCT ( $n=14$ ; pre  $47 \pm 15\%$  vs. post  $46 \pm 19\%$ ; paired  $t$ -test  $P=0.772$ ) or CON ( $n=11$ ; pre  $40 \pm 18\%$  vs. post  $40 \pm 17\%$ ;  $P=0.816$ ).

#### *Intrinsic contractile properties and muscle size*

Both training groups increased Octet Peak T (paired  $t$ -test ECT  $P=0.001$ , SCT  $P=0.015$ ) and Octet TPT (ECT  $P=0.017$ , SCT  $P<0.001$ ), with no change after CON ( $0.689 \leq P \leq 0.986$ ; Table 3). Increases in Octet TPT were greater after SCT than CON ( $ES=1.35$  “large”, LSD  $P=0.009$ ), but not for other comparisons ( $P \geq 0.132$ ,  $0.42 \leq ES \leq 0.74$  “small” to “large”). No changes in Octet T<sub>50</sub> occurred after ECT, SCT or CON (paired  $t$ -test  $0.489 \leq P \leq 0.857$ ), although Relative Octet T<sub>50</sub> decreased after ECT and SCT (both paired  $t$ -test  $P=0.001$ ), but not CON ( $P=0.638$ ; Table 3). There was no ANCOVA interaction effect for Octet T<sub>50</sub> (Table 3), however the decreases in Relative Octet T<sub>50</sub> after both ECT ( $ES=1.36$  “large”, LSD  $P=0.086$ ) and SCT ( $ES=1.37$  “large”,  $P=0.003$ ) were greater than CON, but these changes were similar after ECT and SCT ( $ES=0.25$  “small”,  $P=0.209$ ; Fig. 4).

Twitch Peak T was unchanged in all three groups (paired  $t$ -test  $0.127 \leq P \leq 0.821$ ), although Twitch TPT was longer after both training interventions ( $0.009 \leq P \leq 0.047$ ; Table 3), but not CON ( $P=0.132$ ). No changes in Twitch T<sub>50</sub> occurred after ECT, SCT, or CON (paired  $t$ -test  $0.489 \leq P \leq 0.857$ ). Relative Twitch T<sub>50</sub> decreased after SCT and ECT (paired  $t$ -test  $0.008 \leq P \leq 0.032$ ), but not CON ( $P=0.919$ ; Table 3).

QUADS<sub>VOL</sub> increased 8.1% after SCT from  $1820 \pm 274$  to  $1967 \pm 316$  cm<sup>3</sup> ( $n=15$ ; paired  $t$ -test  $P=0.001$ ), but not following ECT ( $n=13$ ;  $1770 \pm 252$  to  $1816 \pm 286$  cm<sup>3</sup>;  $P=0.247$ ) or CON ( $n=14$ ;  $1891 \pm 272$  to  $1906 \pm 261$  cm<sup>3</sup>;  $P=0.550$ ; Table 4). There was a group x time interaction effect for QUADS<sub>VOL</sub> (ANCOVA,  $P=0.018$ ), with the change in QUADS<sub>VOL</sub> after SCT being greater than that following CON (ES=1.15 “large”, LSD  $P=0.021$ ) and ~3-fold greater than after ECT (ES=0.74 “moderate”,  $P=0.074$ ; Fig. 5). Increases in QUADS<sub>VOL</sub> after ECT were not greater than CON (ES=0.27 “small”, LSD  $P=0.552$ ).

## Discussion

This study compared the specificity of functional adaptations to 12-weeks of ECT vs. SCT and assessed underpinning neural, contractile, and hypertrophic adaptations contributing to these functional changes. MVT increased after both SCT and ECT, but these changes were greater after SCT (+23 vs. +17%). Increases in EMG<sub>MVT</sub> were similar following SCT and ECT, whilst greater increases in QUADS<sub>VOL</sub> (+8.1 vs. +2.6%) suggest muscle size rather than neural drive explained the greater improvement in MVT after SCT than ECT. Improvements in early-phase explosive torque production ( $\leq 100$  ms) only occurred after ECT (+17-34%), were greater than after SCT (at 100 ms) and appeared to be due to increased early-phase neural drive. ECT and SCT both improved explosive strength at 150 ms (+18% vs. +12%) with corresponding increases in neural drive likely explaining the enhancement in late-phase explosive torque production. Octet Peak T increased after training, but there were no changes in the intrinsic contractile explosive capability (Twitch and Octet  $T_{50}$ ) as the time-course of the evoked response (Octet and Twitch TPT as well as Relative Octet and Twitch  $T_{50}$ ) decreased after both SCT and ECT, indicating a likely slowing of the muscle’s contractile properties after both training interventions. Overall, the results support our hypothesis of

distinct and specific functional changes (ECT>SCT for early-phase explosive strength; SCT>ECT for maximum strength), and this appeared to be due to distinct neural and hypertrophic, but not intrinsic contractile adaptations.

Both ECT and SCT increased maximum strength, and by more than CON, but with greater increases after SCT (+23 vs. +17%). Maximum strength has been reported to increase by varying extents following both SCT (+11-36%; (1, 4, 9, 24, 40)) and ECT (+7-25%; (7, 41, 48)), yet this study is the first to directly compare the magnitude of maximum strength improvements after prolonged training with these different approaches. Loading duration (also referred to as time under tension) and loading magnitude have been suggested to be important training stimuli for maximum strength adaptation (15). Maximum strength improvements after ECT were ~70% of those after SCT, despite ECT involving only 7% of the loading duration (time >65%MVT) and thus considerably less effort and fatigue. In contrast the loading magnitude of the two interventions in the current study were physiologically, if not statistically, quite similar (ECT 81 vs. SCT 75%). Overall this provides evidence that loading magnitude rather than loading duration accounts for the majority of the maximum strength improvement following the first 12 weeks of SCT and is the primary training stimulus. In this case, brief explosive contractions up to a high loading magnitude appear to be an efficient means of increasing maximum strength without the requirement for sustained muscular contractions. Furthermore, if loading magnitude is the primary stimulus for maximum strength gains then it is possible that even higher loading magnitudes than those employed in the current study (i.e. >95%MVT), which may be achievable during very short contractions, could provide an even greater stimulus for enhancing maximum strength. The importance of loading magnitude for maximal strength gains may have application for optimizing training prescription of athletes and patient

populations, in particular for patient groups where more sustained contractions may be problematic due to fatigue.

Neural drive at MVT increased more after both SCT and ECT than CON. Numerous previous studies have found neural drive at MVT (assessed with EMG) to increase after SCT interventions (24, 47), however the current study is the first to show that short duration explosive contractions can produce increases in neural drive at MVT and this likely explained the efficacy of ECT for increasing MVT. In fact there was no difference between ECT and SCT for this neural adaptation ( $EMG_{MVT}$ ), indicating that loading magnitude rather than loading duration is the primary stimulus for increasing neural drive at MVT. Previous evidence suggests that increased motor unit firing frequency explains enhanced neural drive at MVT after training (27, 28), and this likely accounts for the improvement of both groups in the current study. In contrast, ECT did not stimulate an increase in muscle volume, and therefore, while ECT appears to be effective at enhancing neural aspects of maximum strength it is relatively ineffective at stimulating hypertrophy. Whereas SCT did induce an increase in muscle volume, that was ~3-fold greater than after ECT (+8.1 vs. +2.6%). Thus hypertrophy was sensitive to loading duration and this adaptation appears to explain the larger improvements in maximum strength for SCT vs. ECT. In this case, for longer-term training goals that are primarily reliant on hypertrophic, rather than neural, adaptations loading duration may become the key training variable. These findings may have relevance for athletic and patient groups where increasing muscle volume is a primary training goal.

Early-phase (first 100 ms) explosive strength increased more after ECT than SCT. In contrast, later-phase explosive strength ( $T_{150}$ ) was enhanced after both types of training. The improvements in  $T_{50}$  and  $T_{100}$  following ECT in the current study are consistent with our

previous observation that early-phase explosive strength adaptations were highly specific to 4-weeks of ECT vs. SCT (45), and demonstrates this to also be the case with more prolonged (12-wks) training. The loading rate (peak RTD) during the short explosive contractions of ECT was almost 6-fold greater than SCT, and therefore, high loading rates, rather than loading magnitudes (similar for ECT and SCT) or duration (greater for SCT) appears to be critical for enhancing early-phase explosive strength. Previous investigations of ECT have consistently reported improvements in explosive strength (7, 22, 23, 41, 45, 48). In contrast, training regimes similar to SCT in the current study have demonstrated both enhanced (1, 6, 9, 13, 29, 44) and unchanged (10, 40, 47) explosive strength. The inconsistent changes in explosive strength in these studies may be partly explained by the variable training instructions provided (e.g. an explosive component (13, 40, 44); no explosive component (6, 9, 47); or unclear (1, 29)). In our laboratory, we have consistently found no increase in early-phase explosive strength after 4 (47) and now 12 weeks of isometric SCT, as well as 3 and 12 weeks of dynamic SCT with isoinertial lifting and lowering (10, 19). Therefore for early-phase explosive strength gains a specific explosive component to the training, involving contractions starting from a low/resting level and performing the rising phase of contraction at a high rate, appears to be important.

Neural drive during the early-phase of explosive contractions increased only after ECT ( $EMG_{0-50}$  and  $EMG_{0-100}$ ; Table 2) and these changes were greater than for CON, but not SCT. The Voluntary  $T_{50}/Octet\ T_{50}$  ratio, which provides an alternate measure of early-phase neural drive, increased from 42 to 53% after ECT, but this was not statistically significant due to the large variability in response between participants. Qualitatively however, the group level Voluntary  $T_{50}/Octet\ T_{50}$  ratio response was notably larger after ECT (+26%) than SCT (-2%) or CON (0%). Later-phase neural drive ( $EMG_{0-150}$ ) was increased after both types of training

(Table 2). Overall, the current study shows neural adaptations during the early-phase of explosive contraction that are specific to ECT, that had previously only been documented for a 4-week training period (45), are still present following a more prolonged intervention. Improvements in early-phase explosive torque production ( $T_{50}$  and  $T_{100}$ ) occurred after ECT without increases in muscle size or early-phase intrinsic contractile capacity for explosive torque production (Octet and Twitch  $T_{50}$ ), supporting the importance of neural drive adaptations for the enhancement of early-phase explosive strength following training.

Explosive torque and EMG expressed relative to corresponding maximum force and EMG were unchanged with ECT but decreased with SCT (Tables 1-2 and Fig. 3 B and D); highlighting further the comprehensive adaptations to ECT (i.e. proportional increases in both explosive and maximum torques and corresponding neural drive) but not SCT (i.e. increases in only maximum torque and neural drive). These changes after ECT partly oppose our previous findings of a greater proportion of maximum strength and EMG being expressed during explosive contractions after 4 weeks of ECT (45), that may be explained by the apparent slowing of the contractile properties and/or greater changes in MVT, and neural drive at MVT, after ECT in the current study. Neurologically, increases in instantaneous motor unit discharge rates and the number of motor units able to produce high discharge rates during explosive contractions and a degree of transfer of these adaptations to maximum contractions may explain the increases in explosive (early- and late-phase) and maximum neural drive after ECT (16, 17). In contrast, the low loading rates ( $385 \text{ Nm.s}^{-1}$ ) but high loading magnitudes (75%MVT) with SCT may have only stimulated adaptations in discharge rate during the production of larger torques (i.e. the late-phase of explosive torque production and the plateau phase of contraction) (27, 28).

Overall, ECT denoted by brief contractions with high RTD produced a wider range of functional adaptations than SCT, with improvements in early- and late-phase explosive strength, as well as maximum strength (Table 4). In contrast, SCT only improved maximum strength and late-phase explosive strength (Table 4). The substantially lower loading duration of ECT (7% of SCT) makes this a less-demanding training modality compared to SCT, which may be preferentially tolerated by musculoskeletal patients and older adults. Future research should investigate: (i) whether ECT may be preferentially tolerated by musculoskeletal patients and older adults, and (ii) also evaluate the efficacy of ECT, and underpinning neuromuscular adaptations, in an isoinertial dynamic training model that is more widely accessible.

The within-group increase in Octet Peak T following both ECT and SCT demonstrated an increase in the maximum contractile capacity of the muscle-tendon unit, although between-group differences were not detected. In contrast, Twitch Peak T was unresponsive to training even after SCT that induced hypertrophic adaptations. Changes in the time-course of evoked responses (Octet and Twitch TPT as well as Relative Octet and Twitch  $T_{50}$ ) indicated an overall slowing of the contractile properties of the muscle tendon unit after both types of training. This apparent slowing of the intrinsic contractile properties is likely due to decreased expression of myosin heavy chain type IIX fibres after training (2, 3, 11). For SCT, the slower contractile properties may explain why during the early-phase of explosive voluntary contraction relative torque decreased and absolute torque remained unchanged, despite increases in maximum strength. After ECT the slower contractile properties may explain why relative explosive torque remained unchanged despite improved neural drive, and why the increases in absolute explosive torque were more modest than might have been expected based on our previous 4-week training study (45) when presumably any potentially



negative morphological changes would have been more limited. Furthermore, even after the brief explosive contractions of ECT the intrinsic contractile properties of the muscle were slowed, which might suggest that these changes may be unavoidable with strength training of previously untrained individuals.

In conclusion, functional, neural, and hypertrophic adaptations showed marked training specificity. ECT produced wide ranging functional adaptations with increases in early- and late-phase explosive and maximum strength due to neural adaptations, and the very low loading duration of ECT (7% of SCT) makes this a substantially less demanding training modality that may be preferentially tolerated by musculoskeletal patients and older adults. SCT produced a greater improvement in maximum strength, but no improvements in early-phase explosive strength. The similar changes in neural drive at MVT after ECT and SCT (despite a lesser gain in MVT following ECT) indicate that this adaptation is largely dependent on loading magnitude. In contrast the ~3-fold greater hypertrophy after SCT than ECT indicates that this adaptation is dependent on loading duration. Improvements in early-phase explosive torque production ( $\leq 100$  ms) appear to rely on a high RTD to induce specific neural adaptations. Finally, an apparent slowing of the intrinsic contractile properties of the muscle-tendon unit after both types of training likely compromises improvements in explosive strength.

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599 **References**

- 600 1. **Aagaard P, Simonsen EB, Andersen JL, Magnusson P, Dyhre-Poulsen P.**  
601 Increased rate of force development and neural drive of human skeletal muscle  
602 following resistance training. *J Appl Physiol* 93: 1318–1326, 2002.
- 603 2. **Adams GR, Hather BM, Baldwin KM, Dudley GA.** Skeletal muscle myosin heavy  
604 chain composition and resistance training. *J Appl Physiol* 74: 911–915, 1993.
- 605 3. **Andersen J, Aagaard P.** Myosin heavy chain IIX overshoot in human skeletal muscle.  
606 *Muscle Nerve* 23: 1095–1104, 2000.
- 607 4. **Andersen LL, Andersen JL, Zebis MK, Aagaard P.** Early and late rate of force  
608 development: Differential adaptive responses to resistance training? *Scand J Med Sci*  
609 *Sport* 20: 162–169, 2010.
- 610 5. **Atkinson G.** Analysis of repeated measurements in physical therapy research: multiple  
611 comparisons amongst level means and multi-factorial designs. *Phys Ther Sport* 3: 191–  
612 203, 2003.
- 613 6. **Balso C Del, Cafarelli E.** Neural Changes Associated with Training Adaptations in  
614 the activation of human skeletal muscle induced by short-term isometric resistance  
615 training. *J Appl Physiol* 103: 402–411, 2007.
- 616 7. **Barry BK, Warman GE, Carson RG.** Age-related differences in rapid muscle  
617 activation after rate of force development training of the elbow flexors. *Exp Brain Res*  
618 162: 122–132, 2005.
- 619 8. **Blazevich A.** Are training velocity and movement pattern important determinants of  
620 muscular rate of force development enhancement? *Eur J Appl Physiol* 112: 3689–91,  
621 2012.
- 622 9. **Blazevich AJ, Horne S, Cannavan D, Coleman DR, Aagaard P.** Effect of  
623 contraction mode of slow-speed resistance training on the maximum rate of force  
624 development in the human quadriceps. *Muscle and Nerve* 38: 1133–1146, 2008.
- 625 10. **Buckthorpe M, Erskine RM, Fletcher G, Folland JP.** Task-specific neural  
626 adaptations to isoinertial resistance training. *Scand J Med Sci Sport* 25: 1–10, 2014.
- 627 11. **Carroll TJ, Abernethy PJ, Logan P a., Barber M, McEniery MT.** Resistance  
628 training frequency: Strength and myosin heavy chain responses to two and three bouts  
629 per week. *Eur J Appl Physiol Occup Physiol* 78: 270–275, 1998.
- 630 12. **Carroll TJ, Riek S, Carson RG.** Corticospinal responses to motor training revealed  
631 by transcranial magnetic stimulation. [Online]. *Exerc Sport Sci Rev* 29: 54–9, 2001.  
632 <http://www.ncbi.nlm.nih.gov/pubmed/11337823>.
- 633 13. **Caserotti P, Aagaard P, Buttrup Larsen J, Puggaard L.** Explosive heavy-resistance

- 634 training in old and very old adults: Changes in rapid muscle force, strength and power.  
635 *Scand J Med Sci Sport* 18: 773–782, 2008.
- 636 14. **Craig CL, Marshall AL, Sjöström M, Bauman AE, Booth ML, Ainsworth BE,**  
637 **Pratt M, Ekelund U, Yngve A, Sallis JF, Oja P.** International physical activity  
638 questionnaire: 12-Country reliability and validity. *Med Sci Sports Exerc* 35: 1381–  
639 1395, 2003.
- 640 15. **Crewther B, Keogh J, Cronin J, Cook C.** Possible stimuli for strength and power  
641 adaptation: Acute Hormonal Responses. *Sport Med* 36: 215–238, 2006.
- 642 16. **Van Cutsem M, Duchateau J, Hainaut K.** Changes in single motor unit behaviour  
643 contribute to the increase in contraction speed after dynamic training in humans. *J*  
644 *Physiol* 513 ( Pt 1: 295–305, 1998.
- 645 17. **Duchateau J, Baudry S.** Maximal discharge rate of motor units determines the  
646 maximal rate of force development during ballistic contractions in human. *Front Hum*  
647 *Neurosci* 8: 9–11, 2014.
- 648 18. **Duchateau J, Enoka RM.** Neural adaptations with chronic activity patterns in able-  
649 bodied humans. *Am J Phys Med Rehabil* 81: S17–27, 2002.
- 650 19. **Erschine RM, Fletcher G, Folland JP.** The contribution of muscle hypertrophy to  
651 strength changes following resistance training. *Eur J Appl Physiol* 114: 1239–1249,  
652 2014.
- 653 20. **Folland JP, Buckthorpe MW, Hannah R.** Human capacity for explosive force  
654 production: Neural and contractile determinants. *Scand. J. Med. Sci. Sport.* (2014). doi:  
655 10.1111/sms.12131.
- 656 21. **Gandevia SC.** Spinal and supraspinal factors in human muscle fatigue. *Physiol Rev* 81:  
657 1725–89, 2001.
- 658 22. **Geertsens SS, Lundbye-Jensen J, Nielsen JB.** Increased central facilitation of  
659 antagonist reciprocal inhibition at the onset of dorsiflexion following explosive  
660 strength training. *J Appl Physiol* 105: 915–922, 2008.
- 661 23. **Gruber M, Gruber SBH, Taube W, Schubert M, Beck SC, Gollhofer A.**  
662 Differential effects of ballistic versus sensorimotor training on rate of force  
663 development and neural activation in humans. *J Strength Cond Res* 21: 274–282, 2007.
- 664 24. **Häkkinen K, Newton RU, Gordon SE, McCormick M, Volek JS, Nindl BC,**  
665 **Gotshalk LA, Campbell WW, Evans WJ, Häkkinen A, Humphries BJ, Kraemer**  
666 **WJ.** Changes in muscle morphology, electromyographic activity, and force production  
667 characteristics during progressive strength training in young and older men. *J Gerontol*  
668 *A Biol Sci Med Sci* 53: B415–B423, 1998.
- 669 25. **Hunter GR, McCarthy JP, Bamman MM.** Effects of resistance training on older  
670 adults. *Sports Med* 34: 329–348, 2004.
- 671 26. **Husby VS, Helgerud J, Bjørgen S, Husby OS, Benum P, Hoff J.** Early Maximal  
672 Strength Training Is an Efficient Treatment for Patients Operated With Total Hip  
673 Arthroplasty. *Arch Phys Med Rehabil* 90: 1658–1667, 2009.
- 674 27. **Kamen G, Knight CA.** Training-related adaptations in motor unit discharge rate in  
675 young and older adults. *J Gerontol A Biol Sci Med Sci* 59: 1334–1338, 2004.

- 676 28. **Knight CA, Kamen G.** Relationships between voluntary activation and motor unit  
677 firing rate during maximal voluntary contractions in young and older adults. *Eur J*  
678 *Appl Physiol* 103: 625–630, 2008.
- 679 29. **Kubo K, Kanehisa H, Fukunaga T.** Effects of isometric training on the elasticity of  
680 human tendon structures in vivo. *J Appl Physiol* 91: 26–32, 2001.
- 681 30. **Lakens D.** Calculating and reporting effect sizes to facilitate cumulative science: A  
682 practical primer for t-tests and ANOVAs. *Front Psychol* 4: 1–12, 2013.
- 683 31. **Linnamo V, Häkkinen K, Komi P V.** Neuromuscular fatigue and recovery in  
684 maximal compared to explosive strength loading. *Eur J Appl Physiol Occup Physiol*  
685 77: 176–181, 1998.
- 686 32. **Liu CJ, Lathan NK.** Progressive resistance strength training for improving physical  
687 function in older adults ( Review ). *Cochrane Libr* : 1–272, 2009.
- 688 33. **Maffiuletti NA, Aagaard P, Blazevich AJ, Folland JP, Tillin NA, Duchateau J.**  
689 Rate of force development: physiological and methodological considerations. *Eur J*  
690 *Appl Physiol* 2016, [date unknown].
- 691 34. **McQuade KJ, De Oliveira AS.** Effects of progressive resistance strength training on  
692 knee biomechanics during single leg step-up in persons with mild knee osteoarthritis.  
693 *Clin Biomech* 26: 741–748, 2011.
- 694 35. **Mizner RL, Petterson SC, Stevens JE, Axe MJ, Snyder-Mackler L.** Preoperative  
695 quadriceps strength predicts functional ability one year after total knee arthroplasty. *J*  
696 *Rheumatol* 32: 1533–1539, 2005.
- 697 36. **Overman CL, Kool MB, Silva JAP Da, Geenen R.** The prevalence of severe fatigue  
698 in rheumatic diseases : an international study. (2015). doi: 10.1007/s10067-015-3035-6.
- 699 37. **Penninx BW, Messier SP, Rejeski WJ, Williamson JD, DiBari M, Cavazzini C,**  
700 **Applegate WB, Pahor M.** Physical exercise and the prevention of disability in  
701 activities of daily living in older persons with osteoarthritis. *Arch Intern Med* 161:  
702 2309–2316, 2001.
- 703 38. **Power JD, Badley EM, French MR, Wall AJ, Hawker GA.** Fatigue in osteoarthritis :  
704 a qualitative study. 8: 1–8, 2008.
- 705 39. **Ratamess NA, Alvar B., Evetoch T., Housh TJ, Kibler W., Kraemer WJ, Triplett**  
706 **N.** Progression Models in Resistance Training for Healthy Adults. (2009). doi:  
707 10.1249/MSS.0b013e3181915670.
- 708 40. **Rich C, Cafarelli E.** Submaximal motor unit firing rates after 8 wk of isometric  
709 resistance training. *Med Sci Sports Exerc* 32: 190–196, 2000.
- 710 41. **de Ruiter CJ, Hutter V, Icke C, Groen B, Gemmink A, Smilde H, de Haan A.** The  
711 effects of imagery training on fast isometric knee extensor torque development. *J*  
712 *Sports Sci* 30: 166–174, 2012.
- 713 42. **Scott SM, Hughes AR, Galloway SDR, Hunter AM.** Surface EMG characteristics of  
714 people with multiple sclerosis during static contractions of the knee extensors. (2011).  
715 doi: 10.1111/j.1475-097X.2010.00972.x.
- 716 43. **Stelmach G, Teasdale N, Phillips J, Worringham C.** Force production  
717 characteristics in Parkinson's disease. *Exp Brain Res* 76: 165–172, 1989.

44. **Suetta C, Aagaard P, Rosted A, Jakobsen AK, Duus B, Kjaer M, Magnusson SP.** Training-induced changes in muscle CSA, muscle strength, EMG, and rate of force development in elderly subjects after long-term unilateral disuse. *J Appl Physiol* 97: 1954–1961, 2004.
45. **Tillin NA, Folland JP.** Maximal and explosive strength training elicit distinct neuromuscular adaptations, specific to the training stimulus. *Eur J Appl Physiol* 114: 365–374, 2014.
46. **Tillin NA, Jimenez-Reyes P, Pain MTG, Folland JP.** Neuromuscular performance of explosive power athletes versus untrained individuals. *Med Sci Sports Exerc* 42: 781–790, 2010.
47. **Tillin NA, Pain MTG, Folland JP.** Short-term unilateral resistance training affects the agonist-antagonist but not the force-agonist activation relationship. *Muscle and Nerve* 43: 375–384, 2011.
48. **Tillin NA, Pain MTG, Folland JP.** Short-term training for explosive strength causes neural and mechanical adaptations. *Exp Physiol* 97: 630–41, 2012.
49. **Tillin NA, Pain MTG, Folland JP.** Identification of contraction onset during explosive contractions. Response to Thompson et al. “Consistency of rapid muscle force characteristics: Influence of muscle contraction onset detection methodology” [*J Electromyogr Kinesiol* 2012;22(6):893-900]. *J Electromyogr Kinesiol* 23: 991–994, 2013.

## Figure captions

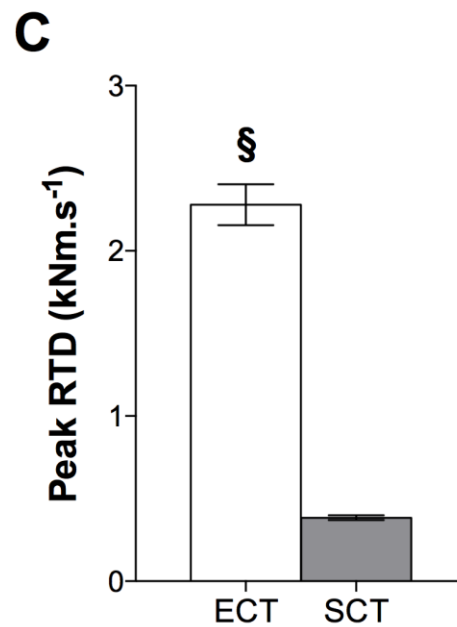
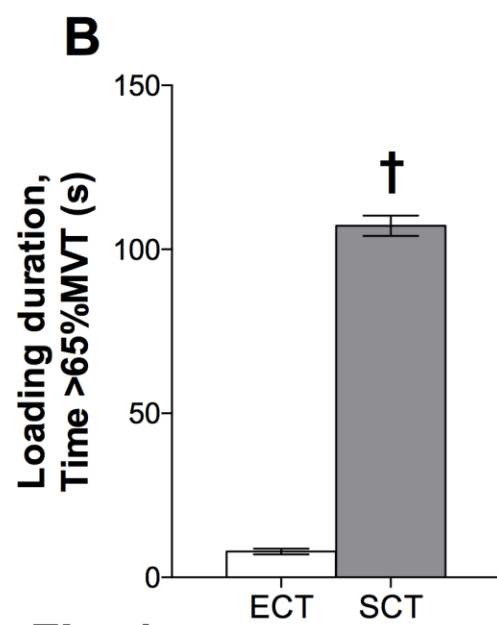
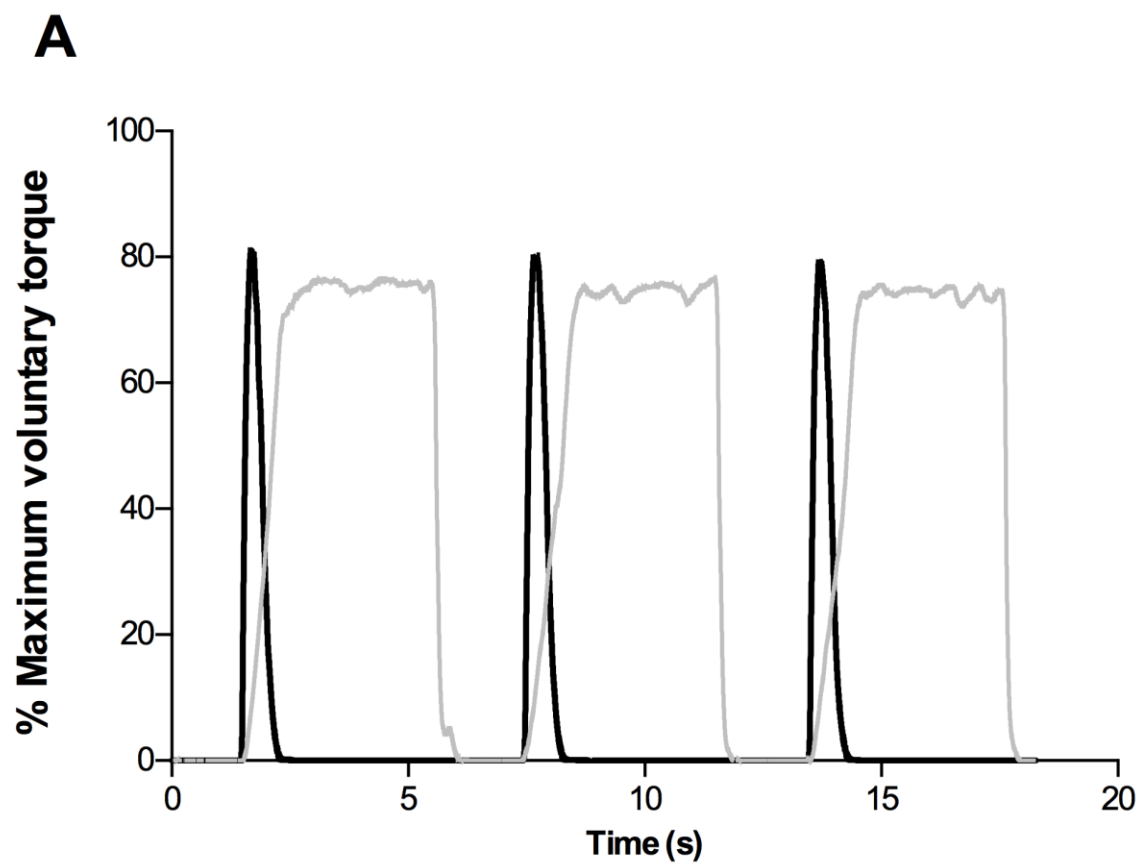
**Fig. 1.** (A) Example torque–time curves recorded during three isometric knee extension contractions for two participants performing either explosive-contraction strength training (ECT; black line) or sustained-contraction strength training (SCT; grey line); (B) Loading duration per training session measured by time >65 maximum voluntary torque (MVT) for ECT vs. SCT; and (C) Peak rate of torque development (RTD, 50 ms epoch) during training contractions for ECT vs. SCT. Symbols indicate differences between training groups as determined from unpaired *t*-tests and are denoted by: †Greater than ECT, §Greater than SCT. Data are mean  $\pm$  SE.

**Fig. 2.** Changes in maximum voluntary torque (MVT) and quadriceps EMG RMS amplitude at MVT (QEMG<sub>MVT</sub>) during isometric knee extensions after explosive-contraction strength training (ECT), sustained-contraction strength training (SCT), and control (CON) interventions. Symbols indicate differences in the magnitude of pre to post changes where post-hoc tests displayed both effect size >0.50 and least significant difference  $P < 0.10$ : \*Greater than CON, †Greater than ECT. Data are mean  $\pm$  SE.

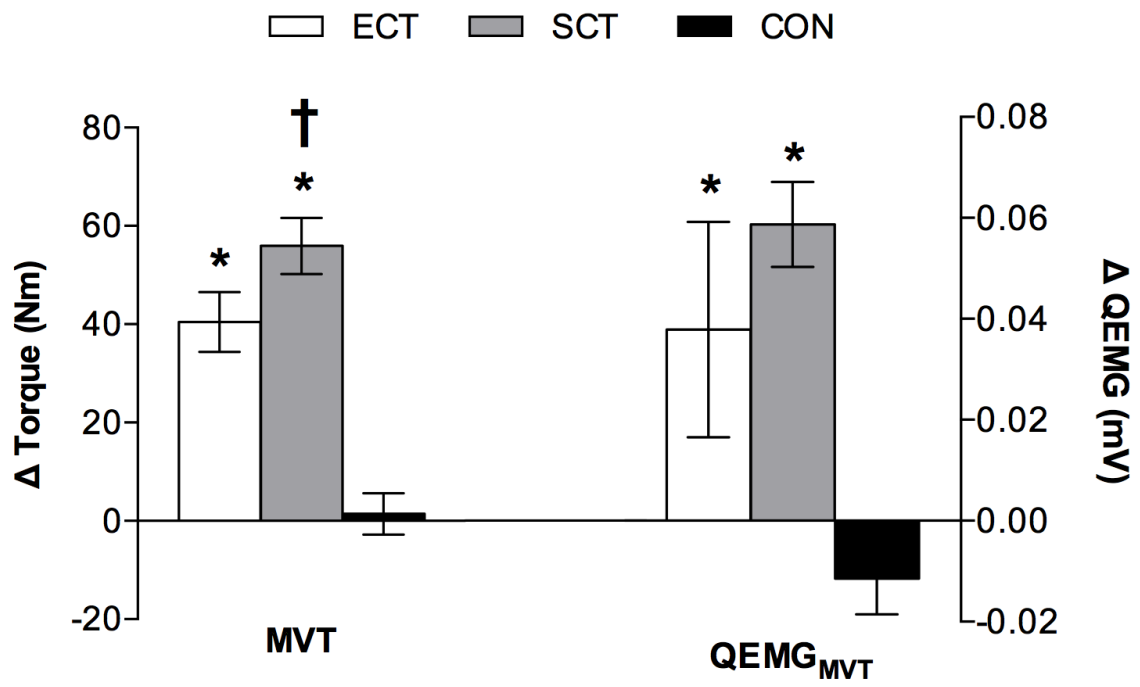
**Fig. 3** Changes in (A) torque, (B) relative torque (%MVT), (C) quadriceps EMG RMS amplitude, and (D) relative explosive quadriceps EMG RMS amplitude (%QEMG<sub>MVT</sub>) during explosive isometric knee extensions after explosive-contraction strength training (ECT), sustained-contraction strength training (SCT), and control (CON) interventions. Symbols indicate differences in the magnitude of pre to post changes where post-hoc tests displayed both effect size >0.50 and least significant difference  $P < 0.10$ : \*Different to CON, †Different to ECT, §Different to SCT. Data are mean  $\pm$  SE.

**Fig. 4.** Pre to post changes in Relative Octet  $T_{50}$  (the ratio between octet torque 50 ms after contraction onset and octet peak torque) after explosive-contraction strength training (ECT,  $n=12$ ), sustained-contraction strength training (SCT,  $n=14$ ), and control (CON,  $n=11$ ) interventions. Symbols indicate differences in the magnitude of pre to post changes where post-hoc tests displayed both effect size >0.50 and least significant difference  $P < 0.10$ : \*Different to CON. Data are mean  $\pm$  SE.

**Fig. 5.** Pre to post changes in total quadriceps muscle volume (QUADS<sub>VOL</sub>) after explosive-contraction strength training (ECT,  $n=13$ ), sustained-contraction strength training (SCT,  $n=15$ ), and control (CON,  $n=14$ ) interventions. Symbols indicate differences in the magnitude of pre to post changes where post-hoc tests displayed both effect size >0.50 and least significant difference  $P < 0.10$ : \*Greater than CON, †Greater than ECT. Data are mean  $\pm$  SE.

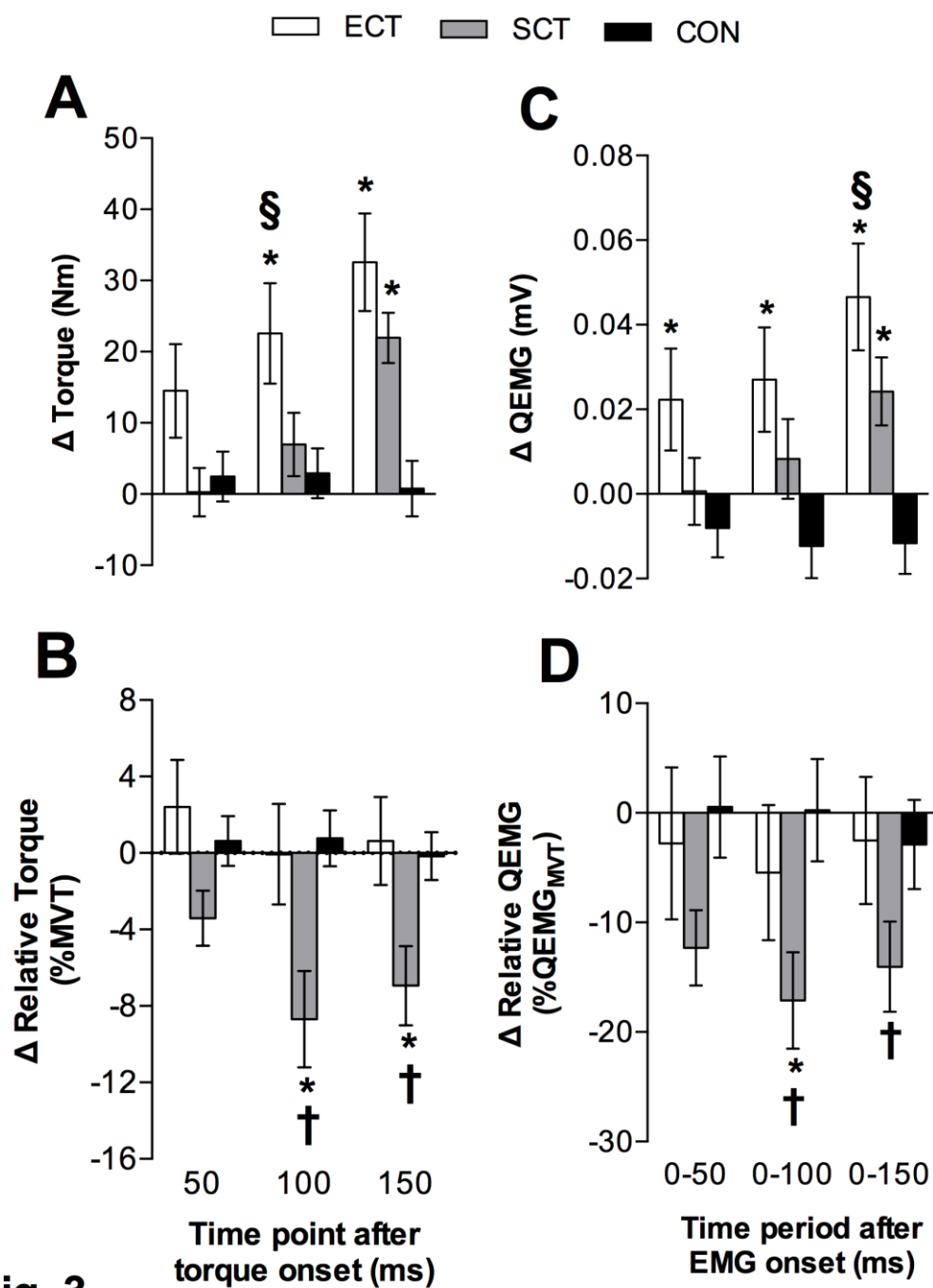


**Fig. 1.**

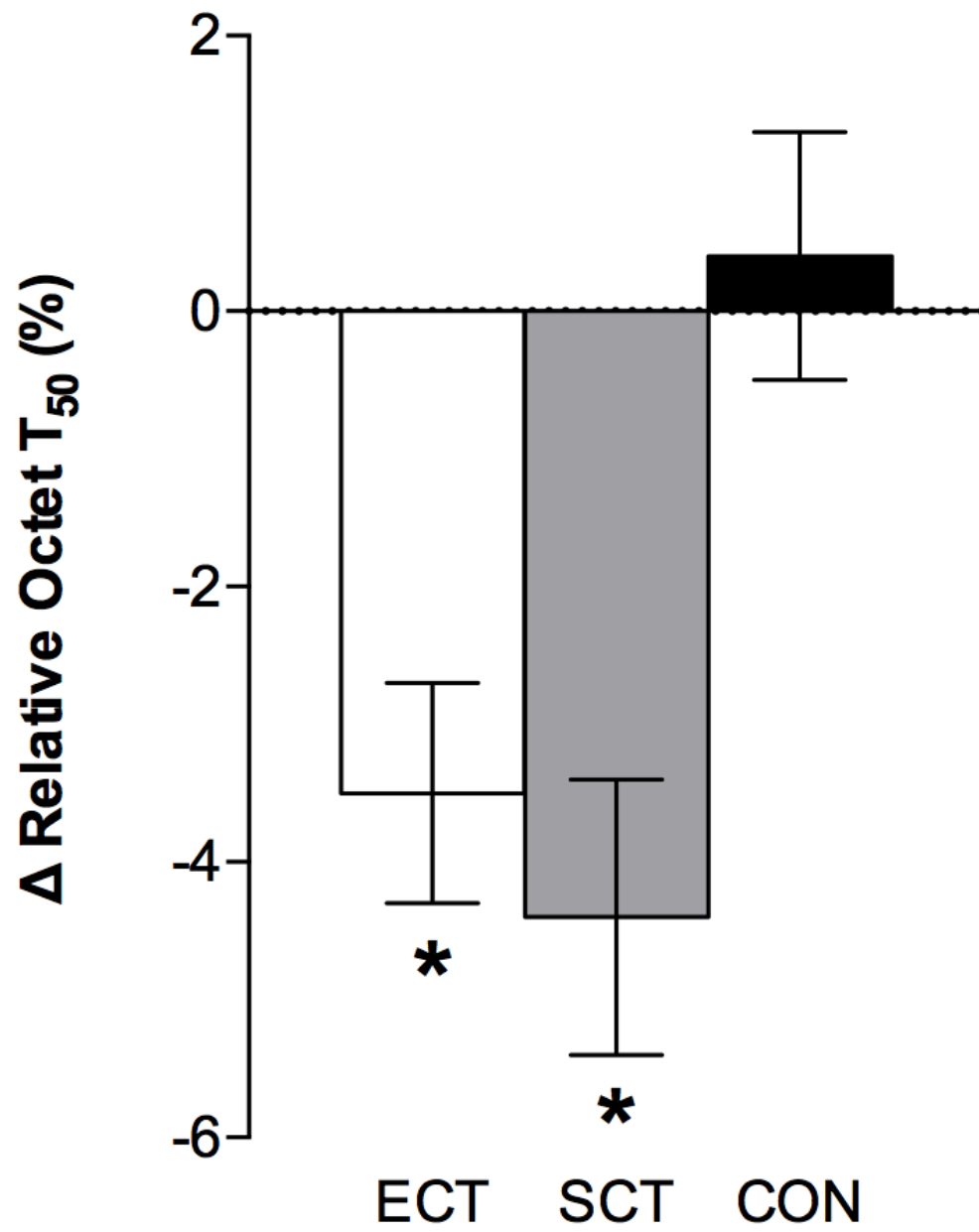


**Fig. 2.**

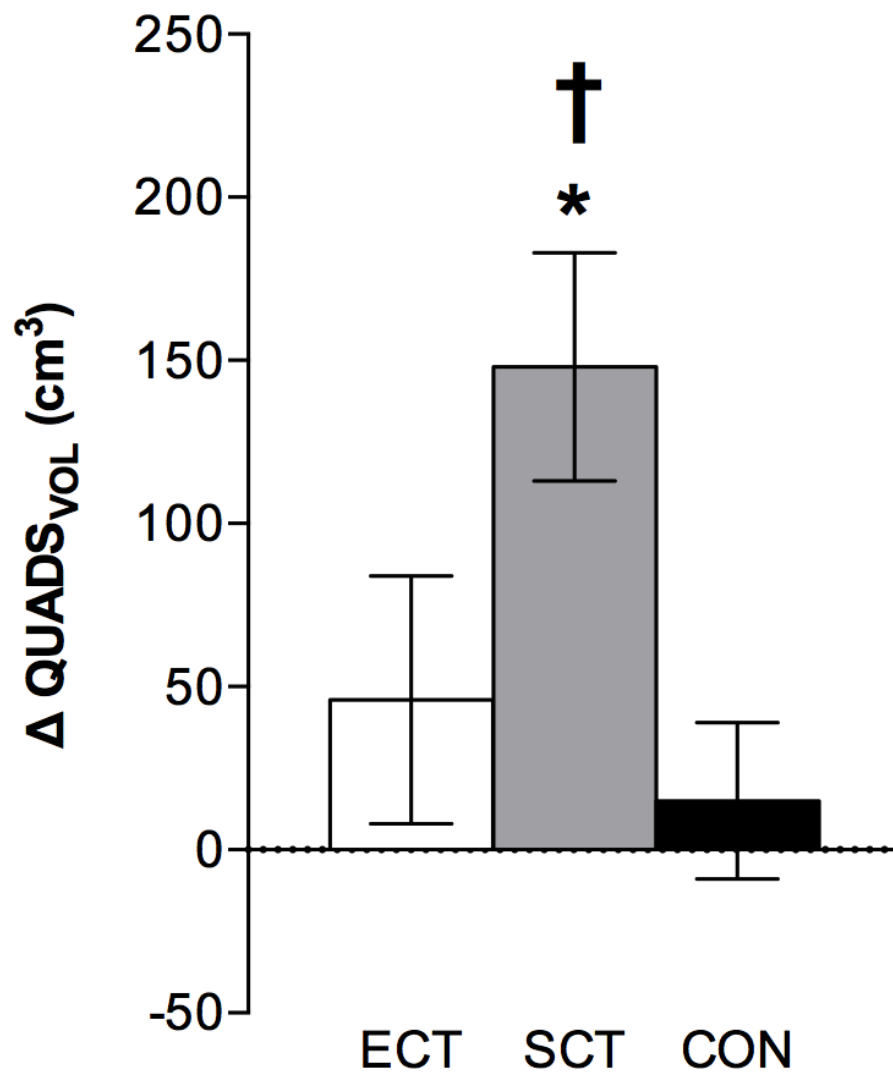




**Fig. 3.**



**Fig. 4.**



**Fig. 5.**

**Table 1.** Maximum voluntary torque (MVT) and explosive torque production (absolute and relative to MVT) pre and post explosive-contraction strength training (ECT,  $n=13$ ), sustained-contraction strength training (SCT,  $n=16$ ), and control (CON,  $n=14$ ) interventions. Explosive torque production is also expressed relative to MVT.

|                         | ECT      |             | SCT      |             | CON      |          | ANCOVA<br>interaction<br>(P value) |
|-------------------------|----------|-------------|----------|-------------|----------|----------|------------------------------------|
|                         | Pre      | Post        | Pre      | Post        | Pre      | Post     |                                    |
| <b>Absolute (Nm):</b>   |          |             |          |             |          |          |                                    |
| MVT                     | 232 ± 27 | 272 ± 37*** | 239 ± 48 | 295 ± 46*** | 257 ± 49 | 259 ± 57 | <0.001                             |
| T <sub>50</sub>         | 43 ± 20  | 57 ± 23*    | 47 ± 21  | 47 ± 19     | 39 ± 19  | 42 ± 19  | 0.058                              |
| T <sub>100</sub>        | 132 ± 25 | 155 ± 29**  | 138 ± 28 | 145 ± 22    | 138 ± 26 | 141 ± 27 | 0.036                              |
| T <sub>150</sub>        | 177± 27  | 210 ± 35*** | 182 ± 34 | 204 ± 25*** | 192 ± 31 | 193 ± 35 | <0.001                             |
| <b>Relative (%MVT):</b> |          |             |          |             |          |          |                                    |
| T <sub>50</sub>         | 18 ± 8   | 21 ± 7      | 20 ± 8   | 16 ± 7*     | 16 ± 7   | 16 ± 6   | 0.055                              |
| T <sub>100</sub>        | 57 ± 8   | 57 ± 7      | 59 ± 10  | 50 ± 7**    | 55 ± 9   | 55 ± 9   | 0.007                              |
| T <sub>150</sub>        | 76 ± 6   | 77 ± 6      | 77 ± 9   | 70 ± 7**    | 75 ± 8   | 75 ± 7   | 0.004                              |

Data are mean  $\pm$  SD. Within-group effects of training were determined from paired *t*-tests and are denoted by: \* (P<0.05), \*\* (P<0.01), or \*\*\* (P<0.001). ANCOVA interaction effects of time (pre vs. post)  $\times$  group (ECT vs. SCT vs. CON) are reported. Post-hoc comparisons of between group changes are shown in Fig. 2 and 3. ECT, explosive-contraction strength training ( $n=13$ ); SCT, sustained-contraction strength training ( $n=16$ ); CON, control; T, explosive torque (at 50 ms intervals from torque onset).

**Table 2.** EMG recorded at maximum voluntary torque (EMG<sub>MVT</sub>) and during explosive contractions (absolute and relative to EMG<sub>MVT</sub>) pre and post explosive-contraction strength training, sustained-contraction strength training, and control interventions.

|                                       | ECT         |               | SCT         |                | CON         |             | ANCOVA<br>interaction<br>(P value) |
|---------------------------------------|-------------|---------------|-------------|----------------|-------------|-------------|------------------------------------|
|                                       | Pre         | Post          | Pre         | Post           | Pre         | Post        |                                    |
| <b>Absolute (mV):</b>                 |             |               |             |                |             |             |                                    |
| EMG <sub>MVT</sub>                    | 0.21 ± 0.08 | 0.25 ± 0.10‡  | 0.18 ± 0.07 | 0.23 ± 0.08*** | 0.19 ± 0.07 | 0.17 ± 0.06 | 0.001                              |
| EMG <sub>0-50</sub>                   | 0.10 ± 0.06 | 0.12 ± 0.07‡  | 0.08 ± 0.05 | 0.08 ± 0.05    | 0.08 ± 0.05 | 0.07 ± 0.04 | 0.033                              |
| EMG <sub>0-100</sub>                  | 0.16 ± 0.07 | 0.18 ± 0.08*  | 0.13 ± 0.05 | 0.13 ± 0.06    | 0.13 ± 0.06 | 0.12 ± 0.05 | 0.022                              |
| EMG <sub>0-150</sub>                  | 0.16 ± 0.07 | 0.21 ± 0.08** | 0.14 ± 0.05 | 0.16 ± 0.06**  | 0.15 ± 0.06 | 0.14 ± 0.05 | <0.001                             |
| <b>Relative (%EMG<sub>MVT</sub>):</b> |             |               |             |                |             |             |                                    |
| EMG <sub>0-50</sub>                   | 49.2 ± 22.8 | 46.5 ± 16.6   | 46.6 ± 21.2 | 34.3 ± 14.4**  | 41.2 ± 17.2 | 41.8 ± 20.6 | 0.102                              |
| EMG <sub>0-100</sub>                  | 78.2 ± 17.6 | 72.7 ± 16.1   | 75.3 ± 23.2 | 58.1 ± 17.3**  | 71.8 ± 16.1 | 72.0 ± 23.6 | 0.031                              |
| EMG <sub>0-150</sub>                  | 83.6 ± 15.9 | 81.1 ± 13.0   | 81.2 ± 19.9 | 67.2 ± 15.9**  | 79.5 ± 15.1 | 76.7 ± 18.2 | 0.048                              |

Data are mean ± SD. Within-group effects of training were determined from paired *t*-tests and are denoted by: \* (P<0.05), \*\* (P<0.01), \*\*\* (P<0.001), or ‡ (P≤0.10). ANCOVA time (pre vs. post) x group (ECT vs. SCT vs. CON) interaction effects are also reported. Post-hoc comparisons of between group changes are shown in Fig. 2 and 3. ECT, explosive-contraction strength training (n =13); SCT, sustained-contraction strength training (n= 16); CON, control; EMG<sub>0-50</sub>, EMG<sub>0-100</sub>, EMG<sub>0-150</sub>, explosive contractions over three time periods from EMG onset (0-50, 0-100, 0-150 ms).

**Table 3.** Intrinsic contractile properties assessed by evoked torque production during octet and twitch contractions pre and post explosive-contraction strength training, sustained-contraction strength training, and control interventions.

|                                     | ECT      |            | SCT      |            | CON      |          | ANCOVA<br>interaction<br>(P value) |
|-------------------------------------|----------|------------|----------|------------|----------|----------|------------------------------------|
|                                     | Pre      | Post       | Pre      | Post       | Pre      | Post     |                                    |
| <b>Octet:</b>                       |          |            |          |            |          |          |                                    |
| Octet T <sub>50</sub> (Nm)          | 101 ± 12 | 105 ± 15   | 107 ± 14 | 106 ± 13   | 108 ± 14 | 109 ± 16 | 0.365                              |
| Octet Peak T (Nm)                   | 159 ± 20 | 174 ± 23** | 171 ± 23 | 183 ± 24*  | 177 ± 26 | 177 ± 26 | 0.077                              |
| Relative Octet T <sub>50</sub> (%)  | 64 ± 5   | 60 ± 4**   | 63 ± 3   | 58 ± 3**   | 61 ± 2   | 61 ± 3   | 0.006                              |
| Octet TPT (ms)                      | 121 ± 7  | 127 ± 7*   | 121 ± 6  | 130 ± 6*** | 123 ± 6  | 124 ± 5  | 0.010                              |
| <b>Twitch:</b>                      |          |            |          |            |          |          |                                    |
| Twitch T <sub>50</sub> (Nm)         | 37 ± 8   | 38 ± 11    | 39 ± 9   | 40 ± 8     | 43 ± 12  | 43 ± 10  | 0.865                              |
| Twitch Peak T (Nm)                  | 43 ± 9   | 45 ± 12    | 47 ± 11  | 50 ± 10    | 52 ± 14  | 52 ± 12  | 0.535                              |
| Relative Twitch T <sub>50</sub> (%) | 86 ± 6   | 83 ± 6*    | 83 ± 5   | 81 ± 4**   | 82 ± 5   | 82 ± 3   | 0.157                              |
| Twitch TPT (ms)                     | 73 ± 8   | 76 ± 7*    | 73 ± 5   | 77 ± 4**   | 78 ± 4   | 76 ± 3   | 0.101                              |

Data are mean ± SD. Within-group effects of training were determined from paired *t* tests and are denoted by: \* (P < 0.05), \*\* (P < 0.01), or \*\*\* (P < 0.001). ANCOVA interaction effects of time (pre vs. post) x group (ECT vs. SCT vs. CON) are reported. Relative octet and twitch measures are expressed as percentage of peak torque during these contractions. Participant numbers for: octet variables, ECT, n=12; SCT, n=14; CON, n=11; twitch variables, ECT, n=13; SCT, n=16; CON, n=14. ECT, explosive-contraction strength training; SCT, sustained-contraction strength training; CON, control.

**Table 4.** Summary of within-group changes from pre to post training in functional, neural, hypertrophic, and intrinsic contractile properties after explosive-contraction strength training (ECT), sustained-contraction strength training (SCT), and control (CON) interventions. The direction of the changes are shown by  $\uparrow$  or  $\downarrow$  with the percentage change in the group mean also shown. Non-significant changes are indicated by  $\leftrightarrow$ .

|   | ECT               | SCT               | CON               |
|---|-------------------|-------------------|-------------------|
| <b>Functional:</b>                      |                   |                   |                   |
| MVT (Nm)                                | $\uparrow+17\%$   | $\uparrow+23\%$   | $\leftrightarrow$ |
| Explosive T <sub>50</sub> (Nm)          | $\uparrow+34\%$   | $\leftrightarrow$ | $\leftrightarrow$ |
| Explosive T <sub>100</sub> (Nm)         | $\uparrow+17\%$   | $\leftrightarrow$ | $\leftrightarrow$ |
| Explosive T <sub>150</sub> (Nm)         | $\uparrow+18\%$   | $\uparrow+12\%$   | $\leftrightarrow$ |
| <b>Neural drive:</b>                    |                   |                   |                   |
| EMG <sub>MVT</sub> (mV)                 | $\uparrow+18\%$   | $\uparrow+33\%$   | $\leftrightarrow$ |
| EMG <sub>0-50</sub> (mV)                | $\uparrow+23\%$   | $\leftrightarrow$ | $\leftrightarrow$ |
| EMG <sub>0-100</sub> (mV)               | $\uparrow+17\%$   | $\leftrightarrow$ | $\leftrightarrow$ |
| EMG <sub>0-150</sub> (mV)               | $\uparrow+28\%$   | $\uparrow+18\%$   | $\leftrightarrow$ |
| <b>Hypertrophy:</b>                     |                   |                   |                   |
| QUADS <sub>VOL</sub> (cm <sup>3</sup> ) | $\leftrightarrow$ | $\uparrow+8\%$    | $\leftrightarrow$ |
| <b>Contractile properties:</b>          |                   |                   |                   |
| Octet Peak T (Nm)                       | $\uparrow+9\%$    | $\uparrow+7\%$    | $\leftrightarrow$ |
| Octet TPT (ms)                          | $\uparrow+5\%$    | $\uparrow+7\%$    | $\leftrightarrow$ |
| Twitch TPT (ms)                         | $\uparrow+4\%$    | $\uparrow+5\%$    | $\leftrightarrow$ |